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(54) HYDROLYZABLE METAL-CONTAINING RESIN AND ANTIFOULING PAINT COMPOSITION

(57) A substrate resin having pendant acid groups is produced by oppolymerizing an acrylic or methacrylic ester of which alcoholic residue includes a bulky hydrocarbon radical or a soft segment. The substrate resin is then metallized together with an organic mobobasic acid so that both of the pendant acid group and the organic monobasic acid group are bount to the same metal atom. The resulting hydrolyzable metal-containing resin is incorporated into self-polishing antifouling paint formulations together with an antifouring pigment such as cuprous oxide.

Description

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BACKGROUND ART

This invention relates to a hydrolyzable, metal-containing resin for use as a vehicle resin of antifouling paints as well as antifouling paint compositions containing said resin.

Antifouling paints containing as a vehicle resin a trialkyltin-containing polymer are known. These resins are advantageous in controlling the release of antifouling agent to a minimum level sufficient to maintain desired antifouling effect at a constant level for a long period of time. In application, the vehicle resin consisting of a trialkyltin-containing polymer is hydrolyzed by the action of weakly alkaline sea water to release the trialkyltin, and at the same time the resin becomes water-soluble so that the paint film is consumed and smoothened. This contributes to the reduction of the frictional resistance of ships against water and, therefore, to the reduction of fuel cost.

The vehicle resins of this type of paints, known as "self-polishing paints", typically consist of copolymers of trialkyltin (meth)acrylate with other comonomers free from carboxyl group. However, strong concern about the toxic effect of trialkyltin on the ecological system has led to a demand for a new vehicle resin which may replace the trialkyltin-containing polymers in the formulation of self-polyshing antifouling paints.

JP-A-62101653, JP-A-62057646, JP-A-63128008 and JP-A-63128084 disclose a metal-containing resin in which a metal atom is ionically combined with an acid pendant group of the resin and further with a monobasic organic acid, and methods of production of such resins. This type of resins may also be hydrolyzed gradually in the sea water to release antifouling metal ions and become soluble by themselves so as to achieve self-polishing effects.

The metal-containing resins are produced by copolymerizing an unsaturated organic acid monomer with a neutral monomer to prepare an acid group-containing resin (hereinafter referred to as "substrate resin"), and then combining the metal and monobasic organic acid. However, the effects of neutral monomers present in the substrate resin on the film performance of antifouling paint formulations containing metallized substrate resins and antifouling pigments have not been investigated well until now. This is because the dissolution rate of films, for example, has long been believed to be affected mainly by the acid number of the subsrate resin but not with the nature of neutral monomers significantly.

DISCLOSURE OF THE INVENTION

We have found that a substrate resin containing certain types of neutral monomers may exhibit, when compared with the corresponding substrate resin not containing such a neutral monomer, enhanced film performance including anti-cracking, adhesive, self-polishing and other properties after metallizing and formulating into an antifouling paint with an antifouling pigment.

The present invention provides a hydrolyzable, metal-containing resin comprising a polymer having in the molecule thereof a plurality of pendant acid groups bound ionically to a metal atom and a monobasic organic acid also bound to the same metal atom as said pendant acid group. It also provides an antifouling paint composition comprising said metal-containing resin and an antifouling pigment and/or antifouling agent. The metal-containing resin according to the present invention is comprised of, in its free acid (substrate resin) form, a copolymer consisting essentially of

- (a) from 5 to 70 % by weight of a member selected from the group consisting of a (meth)acrylic acid ester having as the ester residue a branched alkyl of four or more carbon atoms having at least one branch on a carbon atom at second to fourth positions from the distal end of the principal chain, a (meth)acrylic acid ester having as the ester residue a cycloalkyl residue having six or more carbon atoms, a polyalkylene glycol mono(meth)acrylate, a polyoxyalkylene glycol monoalkyl ether mono(meth)acrylate, and an adduct of 2-hydroxyethyl (meth)acrylate with caprolactone;
- (b) a proportion of a polymerizable unsaturated organic acid monomer corresponding to an acid number of the resin from 25 to 350 mg KOH/g as solid; and
- (c) the balance of another polymerizable neutral monomer.

PREFERRED EMBODIMENTS

The hydrolyzable, metal-containing resin of the present invention may be produced by the methods disclosed in the above-cited patent applications except that the monomeric composition of the substrate resin additionally contains a neutral monomer as defined herein. The pendant acid group e.g. -COOH group binds ionically to a transitional metal, e.g. copper and a monobasic acid, e.g. acetic acid to form the following salt:

-COOCuOAc

The substrate resin having pendant acid groups may be produced by copolymerizing the above monomer (a) with the monomer (b) having an acid group and the monomer (c) free of the acid group in accordance with the conventional solution polymerization method.

Examples of carboxycolacid monomers are adylic and methadrylic acids. Other examples of carboxylic group-containing monomers, notified monoalky, maleate and monoalky stadonate as well as nalf esters of dicarboxylic acid such as pritral of succinic or maleic acid with 2-hydroxyalky/ (meth)acry ate.

Examples of sulfonic group-containing monomer: include pistyrenesulfonic acid. 2-methy-2-acily amdopse-pares of onic acid and the like

Examples of phosphoriloigi bupidantaining monomers include acid phosphorepy, methacrylate, 3-onioro-2-acid phosphory methacrylate, acid phosphoryethy, methacrylate and the like

The manametric composition of the substrate resin should contain the acid gloup-containing monomer in a proportion occidespording to an acid number of the resulting copplymen of from 25 to 350 mg kiOHig as solid. An acid number within this range may generally be reached by compounding the acid matriciper. In a proportion of from 5-76 % preferably from 10 to 50 % by weight of the entire monaments composition. When the proportion of acid monomer lies within the above range, the balance between the durability and the dissolution rate (sembolishing rate, of paint films may be optimized.

In order to impart the paint films with improved adhes on lanti-pracking and other properties, a (meth) adrylic ester monomer cumaring a bulky group or soft segment in the ester residue is copolymenzed. Typical examples of bulky monomers are tibuty adrylate and tibutyl methal, ylate. Other examples of bulky monomers include (meth) adrylic adid esters having as etser residue a branching alky of four time eiga bon atoms having at least one branching a darbon atom at second to fourth positions from the distallend of the principal chemisuith as isobory? Indepenty, ineopenty, or some darbon atoms such as cyclonery) or isobornyl (meth) adrylate.

Folyoxyalkylene chains produced by the ring-opening polymerization of an alkylene oxide such as ethylene oxide or propylene oxide are known to be a soft segment. Examples of (meth)acrylate and desters containing as soft segment a polycizvalkylene chain include polycizvalkylene glycol monormeth)acrylate and polycizvalkylene glycol monormethyl ether methacrylate. Polyoxyethylene (n=8) glycol monormethyl ether methacrylate are dominercially available from Shin Nakamura hagakulkikiki as NK ESTER M-90G and NR ESTER M-2005, respectively. Another example of soft segment-bontaining acrylic monomers is an adduct of 2-hydroxyethyli (methiacrylate with caprolactone. These monomers are also commercially available from Daicel Chemical Industries. Ltd. as FLACCEL FA and PLACCEL FM series.

Medium and long thain akky (methiadrylates are not usable alone because of their high hydrophobidity but usable in combination with a hydrophobid soft segment-centaining adrylic monomer such as NR ESTER M-90G or NR ESTER M-230 3.

The substrate reconshould contain in its monomer occumposition 5-70.7% preferably 10-50.% by weight of the entire monomer is composition of the above monomer for improving adnessin and anti-orational properties of paint films. When the proportion of this kind of monomers lies within the above range it is possible for the metal idea paint films to impart with improved achiesion and durability as well as a constant dissolution rate in the presence of an antifouling big-ment without affecting other properties.

The balance of monomeric composition of the substrate resin is occupied by a neutral monomer other than the monomers (a) and (b). Examples thereof include hydrocarbon monomers such as employed propylene styrene amethylotyrene and unvitoluene alkylometh)acrylates ruch as methylometh)acrylate ethylometh)acrylate hydroxylate such as 2-hydroxyethy (meth)acrylate and hydroxyproxylometh acrylate amides such as acrylamide and methacrylate numbers acrylamide extensionally protected and methacryland extensionally acreated and methacryland extensionally acreated and methacryland extensionally protected.

The substrate reskripreferably has a number average no edular weight from 2,000 to 100,000, more preferably from 3,000 to 40,000. The substrate reskrips reduced to have a more duar weight of the spoke range to haintain an optimal balance almong the fini-forming property, wo kapplity and dissolution rate.

Meta's forming a sait with the substrate resin are prosen from elements of grd ups 3A to 7A 8, 1B and 2B of the period chart. Copa ti thokel, coppor zinc ite killium and manganese are preferable among others.

Polymeric metal saits may be produced by relating the substrate resin with at least a stoichiometric amount of a metal compound such as oxide, hydroxide, phloride is liftue or basic carbonate and at least a stoichiometric amount of a monopasic regard acid simultaneously. Alternative is the sidistrate resin may be leaded with a metal sait of monopasic page.

Examples of monopasion or rand across used to this purpose indicate monoparticity in across such as a certificities of buty of four classes. The order of appropriate form such as a person of phenotypes as a representation of the such as person of person of persons as persons of persons

A preferred method for product of the dolyment metal self is dispresed in . If A 63129008 streams a metalor is A continuity method preductivities in a reamble with a metal self of which industry membrase self-and a five in more open or more self-and a similar terms, to take a power of left data in the first the residuence and and another out of more mateloaded in the self-and and a self-and the round of more mateloaded.

ing point-monobasic acids" as used herein reffered to the existence of substantial difference in boiling points therebetween, for example 20 °C.

The hydrolyzable, metal containing resin thus produced may be incorporated into antifouling paint formulations in combination with an antifouling agent and other conventional additives to prepare the self-polishing antifouling paint composition of the present invention.

The antifouling paint composition according to the present invention may contain the following conventional additives.

(1) Antifouling agent:

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Powders or flakes of copper, zinc and nickel; oxides, halides or hydroxides of copper, zinc and lead; organotin compounds such as tributyltin fluoride or triphenyltin chloride; biocidal metal carboxylates such as copper naphthenate or copper stearate; metal (e.g. Na, K, Zn, Pb, Cu, Fe, Ni, Mg, Se) dialkyl dithiocarbamates such as zinc dimethyl dithiocarbamate and thiuram disulfide; sulfamides such as phthalylsulfathiazole, sulfaethydole, sulfanilidopyridine or sulfamethoxyine; antibiotics such as penicillin V, penicillin G, ampicillin, cephalosporin, chlortetracycline, neomycin, rifamycin or variotin; pyrrole and imidazole compounds such as glyodine, fentizole or polycide; thioxane and thioxanthone compounds such as terazol, asterol or mylone; amides such as nicarbazin, 3,4,5-tribromosalicylanilide, N-trichloromethylmercaptophthalimide or 3,5-dinitrobenzamide; and other known antifouling agents, pesticides, bacteriocides and fungicides.

(2) Plasticizer:

Phthalate plasticizers such as dioctyl phthalate, dimethyl phthalate or dicyclohexyl phthalate; aliphatic dicarboxylate plasticizers such as butyl sebacate; glycol ester plasticizers such as diethylene glycol dibenzoate or pentaerythritol alkanoic etser; phosphate plasticizers such as tricresyl phosphate or trichloroethyl phosphate; epoxy plasticizers such as epoxydized soybean oil or epoxydized octyl stearate; organotin plasticizers such as dioctyltin dilaurate or dibutyltin dilaurate; and other plasticizers such as trioctyl trimellitate, camphor or triacetin.

(3) Hydrolysis regulator:

Chlorinated paraffin, polyvinyl ether, polypropylene sebacate, partially hydrogenated terphenyl, polyvinyl acetate, polyalkyl (meth)acrylate, polyether polyol, alkyd resin, polyester resin and polyvinyl chloride.

(4) Pigment:

Extender pigments such as baryte, precipitated barium sulfate, talc, clay, chalk, silica white, alumina white, titanium white or bentonite; color pigments such as titanium dioxide, zirconium dioxide, basic lead sulfate, tin oxide, carbon black, graphite, red iron oxide, chromium green, emerald green or phthalocyanine blue.

(5) Solvent

Hydrocarbons such as xylene, toluene, benzene, ethylbenzene, cyclopentane, octane, heptane, cyclohexane or white spirit, ethers such as dioxane, tetrahydrofuran, ethylene glycol monomethyl ether, ethylene glycol monomethyl ether, ethylene glycol dibutyl ether, diethylene glycol monomethyl ether or diethylene glycol monomethyl ether; esters such as butyl acetate, propyl acetate, benzyl acetate, ethylene glycol monomethyl ether acetate or ethylene glycol monoethyl ether acetate; ketones such as methyl isobutyl ketone or ethyl isobutyl ketone; and alcohols such as n-butanol or propyl alcohol.

(6) Viscosity regulator:

Amides and amines such as nicotinamide or n-octylamine; monobasic organic acids such as acetic, oleic or lauric acid; phosphoric acid and phosphate esters; solid acids such as silicate or molybdate.

(7) Other additives:

Monobasic organic acids such as rosin, monobutyl phthalate or monooctyl succinate; camphor and castor oil. The antifouling paint composition of the present invention may be prepared by the method known per se in the art. Any known machine such as ball mills, pebble mills, roll mills or spead run mills may be used for mixing various ingredients.

It is desirable for the antibuting paint of the present invention that the hydroryzable resin occurs as a salt with a metal having prization tendency ower than that of alkal metals leight the zono copper or terulium salt. After such a salt is prosen, the paint if insiapplied on ships from retsip manine construction will be gradually nyttiplyzed and dissolved out into weakly alkaline sea water. Opposite to polyester venicle resins containing a number of metal ester molectes in the polymer banknown the venicle lesin of the present invention are not decomposed rapidly into a larger number of soluble fragments but only bendant groups thereof ale hydrolyzed into a hydrophic gloup before the concentration or density of such hydrophic groups reach at a threshold level at which the paint timp begins to dissolve in the sea water. Accordingly, when said vehicle resin is used it is possible to give an antifouring paint timm exhibiting the antifouring and other performance for along period of time. Therefore, the antifouring paint composition according to the present invention finds use in finishing not only show including tankers, ferry boats, steel boats, wood boats and FRP toats but also various manine construction and fish nets.

Besides, particular advantages are appreved when the hydrolyzable, metal-containing resin of the present invention is combiner, with a copper-based antifouring pigment such as outrous oxide or copper inpadance. This is because the interaction between the vehicle resin and the antifouring pigment is remarkably retailed when compared to the corresponding resin not Containing a neutral monomer as defined herein. As a result, the film performance including integrity, superson shength and self-polishing property may be significantly improved.

The invertion is further illustrated by the following production examples, examples and complarative examples in which all parts are by weight.

Froduction Example 1 (Varnish)

To a four necked flask equipped with a stirrer lief us condense, introgen gas tube and drip funnel were added 64 parts of riviene and 16 parts of in-butanot. After heating the content to 100 C, a mixture of 58.3 parts of ethy) acrylate 15 parts of systemesty, methacrylate 10 parts of NK ESTER M-90G (methoxypolyethylene glycol methacrylate sold by Shin Nakaniura Kagaku K.K.), 16.7 parts of acrylic acid and 2 parts of t-butyleleroxy 2-ethylhexanoate was added dropwise at a constant rate over 3 hours. After the addition, the mixture was kept at the same temperature for 30 minutes. Then, a solution of 0.2 parts t-butyleleroxy 2-ethylhexanoate in 16 parts of xylene and 4 parts of in-butanot was added dropwise at a constant rate over 36 minutes. Thereafter the reaction mixture was kept at the same temperature for 1.5 hours. A resin scrution called Vainish A having 49.3 % solids, a viscosity of 4.4 poise and an acid number of 13C (as solids, the same hereinafter) was obtained.

Production Example 2 (Valinish

A mixture of 50 parts of xylene and 50 parts of n-butand was heated to 90. O in the same flask as used in Production Example 1. To this was added drop wise a mixture of 54 6 parts of ethy lastly ate. 7.4 parts of methyl methacilylate 16.7 parts of addyl cladid. 21.8 parts of tibutyl methacilylate and 2 parts of azobis subutylonitrile at a constant rate over 3 hours. After the addition, the mixture was kept at the same temperature for 2 hours. A resin solution called Varinsh B having 50.1.% solds, a viscosity of 5.2 poise and a resin acid number of 130 was obtained.

Production Example 3 Carnish

A mixture of 40 parts of xylene and 40 parts of inputarior was heated to 100.0 in this same flask as used in Production Evample 1. To this was added dropwise a mixture of 65.7 parts of ethy laciylate i 15 parts of cyclohexyl methator, at 6.19.3 parts of acrylolated and 2 parts of tiputy per avected, nevalicate at a constant rate over 3 hours. After the addition the mixture was kept at the same temperature for 30 minutes. Then also ut on 10 parts of 4 parts of tiputy per avectors was added dropwise and constant rate over 30 minutes. Thereafter the reaction mixture was kept at the same temperature for 1.5 hours. A resin solution called Varinsh C having 50.0 %, ocide is also est, of 6.5 poise and a resin acid number of 150 was entained.

-Production Example 4 Lain shi

A mature of 64 parts of 4, whe and the farm of notices was near atom \$7. In the competest as used notices of the parts of some, and the formation of the same of the parts of some, methan parts 220 parts of methan parts 220 parts of methan parts 220 parts of methan parts at the methan parts of the methan of a parts of methan of the formation of the formatio

Production Example 5 (Varnish)

A mixture of 64 parts of xylene and 16 parts of n-butanol was heated to 100°C in the same flask as used in Production Example 1. To this was added dropwise a mixture of 10.2 parts of 2-ethylhexyl methacrylate, 66.8 parts of ethyl acrylate, 11.4 parts of NK ESTER M-90G, 11.6 parts of acrylic acid and 2 parts of t-butylperoxy 2-ethylhexanoate at a constant rate over 3 hours. After the addition, the mixture was kept at the same temperature for 30 minutes. Then a solution of 0.2 parts of t-butyl 2-ethylhexanoate in 16 parts of xylene and 4 parts of n-butanol was added dropwise at a constant rate over 30 minutes. Thereafter the reaction mixture was kept at the same temperature for 1.5 hours. A resin solution called Varnish E having 50.3 % solids, a viscosity of 4.7 poise and a resin acid number of 90 was obtained.

Production Example 6 (Varnish)

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A mixture of 64 parts of xylene and 16 parts of n-butanol was heated to 100°C in the same flask as used in Production Example 1. To this was added dropwise a mixture of 20.0 parts of isobutyl acrylate, 50.9 parts of ethyl acrylate, 14.1 parts of acrylic acid, 15.0 parts of NK ESTER M-90G and 3 parts of t-butylperoxy 2-ethylhexanoate at a constant rate over 4 hours. After the addition, the mixture was kept at the same temperature for 30 minutes. Then a solution of 0.2 parts of t-butylperoxy 2-ethylhexanoate in 16 parts of xylene and 4 parts of n-butanole was added dropwise at a constant rate over 30 minutes. Thereafter the reaction mixture was kept at the same temperature for 2 hours. A resin solution called Varnish F having 50.0 % solids, a viscosity of 7.5 poise and a resin acid number of 110 was obtained.

Production Example 7 (Varnish for comparison)

A mixture of 64 parts of xylene and 16 parts of n-butanol was heated to 100°C. To this was added dropwise a mixture of 54.1 parts of ethyl acrylate, 26.6 parts of methyl methacrylate, 19.3 parts of acrylic acid and 3 parts of azobisisobutyronitrile at a constant rate over 4 hours. After the addition, the mixture was kept at the same temperature for 30 minutes. Then a solution of 0.2 parts of azobisisobutyronitrile in 16 parts of xylene and 4 parts of n-butanol was added dropwise at a constant rate over 30 minutes. Thereafter the reaction mixture was kept at the same temperature for 1.5 hours. A resin solution called Varnish G having 50.5 % solids, a viscosity of 4.5 poise and a resin acid number of 150 was obtained.

Production Example 8 (Varnish for comparison)

A mixture of 50 parts of xylene and 50 parts of n-butanol was heated to 100°C in the same flask as used in Production Example 1. To this ws added dropwise a mixture of 45.6 parts of ethyl acrylate, 32.5 parts of methyl methacrylate, 10.3 parts of 2-hydroxyethyl acrylate, 11.6 parts of acrylic acid and 1.5 parts of t-butylperoxy 2-ethylhexanoate at a constant rate over 3 hours. After the addition, the reaction mixture was kept at the same temperature for 2 hours. A resin solution called Varnish H having 49.8 % solids, a viscosity of 12.9 poise and a resin acid number of 90 was obtained.

40 Example 1

A four necked flask equipped with a stirrer, nitrogen gas tube, reflux condenser, decanter and temperature control means was charged with 100 parts of Varnish A, 25.4 parts of zinc acetate, 24.4 parts of SA-13 (mixture of aromatic sulfonic acids sold by Idemitsu Petrochemical Co., Ltd.) and 140 parts of xylene. The mixture was heated at 130 °C while distilling off acetic acid produced as a by-product with solvent. The end point of the reaction was confirmed by determining the quantity of acetic acid in the effluent solvent. A varnish having 40.2 % solids and a viscosity of U-V was obtained.

Example 2

The same flask as used in Example 1 was charged with 100 parts of Varnish B, 18.6 parts of copper oxalate, 32.7 parts of oleic acid and 120 parts of xylene. The mixture was heated at 120 °C while distilling off oxalic acid with solvent. A varnish having 42.5 % solids and a viscosity of Z_1 - Z_2 was obtained.

55 Example 3

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The same flask as used in Example 1 was charged with 100 parts of Varnish C, 21.5 parts of copper oxalate, 37.5 parts of naphthenoic acid and 120 parts of xylene. The mixture was heated at 120 °C while distilling off oxalic acid with solvent. A varnish having 31.7 % solids and a viscosity of Y-Z was obtained.

Example 4

The same frask as used in Example 1 was charged with 100 parts of Varnish A, 24 0 parts of copper acetate, 30 4 sparts of naphthenologic diand 100 parts of kylene. The multiple was heated at 130 10 while distring off acet cland with solvent. A varnish having 30 5 % soldcland a viscosity of r was soldclaned.

Example 5

The same flask as used in Example 1 was charged with 100 parts of Varnish Di 29.3 parts of zinc acetate, 37.8 parts of bield acid and 120 parts of xylene. The mixture was heated at 130.10 white distilling off acetic acid with solvent. A varnish having 41.3 % solids and a viscosity of VFW was obtained.

Example 6

The same flask as used in Example 1 was charged with 100 parts of Varnish D; 26 0 parts of teliginum abetate, 32 7 parts of Versatid abid and 150 parts of xylene. The mixture was heated at 130. O while distribing off acetic acid with solvent. A carriesh having 55 6 0s solute and a viscosity of S-T was obtained.

Example 7

The same flask as used in Example 1 was charged with 100 parts of Varnish B, 24.0 parts of copper acetate, 32.5 parts of naphthenoic acid and 100 parts of xylene. The mixture was heated at 130. C while distilling off acetic acid with solvent. A varnish having 40.6 % solids and a viscosity of Z--Z, was obtained

JA Example 8

The same flask as used in Example 1 was charged with 100 parts of Varnish E. 31.6 parts of zinc salicylate. 16.9 parts of SA 13 (Idemitsu Petrochemical) and 150 parts of xylene. The mixture was heated at 12010 while distilling off salicylip acid with solvent. A varnish having 62.3 % solids and a viscosity of V-W was obtained.

Example 9

The same flask as used in Example 1 was charged with 100 parts of Varnish C. 26.7 parts of copper acetate i22.7 parts of Versaho acid and 100 parts of kylene. The mixture was heated at 130. C while distribing off acetic acid with solvent. A varnish having 33.4.% so ds and a viscosity of 2.7, was obtained.

Example 10

The same flask as used in Example 1 was charged with 100 parts of Namish Ell16 0 parts of copper acetate 22.5 parts of naphthenoic acid and 120 parts of kylene. The diskture was heated at 130. Own leid stilling off abetic acid with soluent Alvarinsh having 60.7% solids and alviscosity of TIU was obtained.

France

The same flask as used in Example 1 was charged with 100 parts of Camish Fill 1 parts of corpe localate 2010 parts of the alacid and 100 parts of where The mixture was heated at 100 in white disting official class and a viscosity of Wilk was obtained.

A variesh having 50.8 % solids and a viscosity of Wilk was obtained.

Example 10

The same flack as used in Example 1 was charged with 120 be its of Variosh Filiph (practs of the unit masset) as the distribution of the total ordinates of the masset of the description of the masset of the masset of the description of the masset of the m

📑 mgrahat 🕫 Example 1

The land site is used in Example 1 was unarged with the parts of Lambor (B) (A) parts of oppositions as a safety of namenous and parts of case of the constitution of

Comparative Example 2

The same flask as used in Example 1 was charged with 100 parts of Varnish G, 52.6 parts of zinc salicylate, 28.2 parts of SA-13 (Idemitsu Petrochemical) and 150 parts of xylene. The mixture was heated at 120°C while distilling off salicylic acid with solvent. A varnish having 37.4 % solids and a viscosity of W-X was obtained.

Comparative Example 3

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The same flask as used in Example 1 was charged with 100 parts of Varnish H, 16.0 parts of copper acetate, 22.7 parts of oleic acid and 100 parts of xylene. The mixture was heated at 130 °C while distilling off acetic acid with solvent. A varnish having 53.7 % solids and a viscosity of W was obtained.

Comparative Example 4

The same flask as used in Example 1 was charged with 100 parts of Varnish H, 12.9 parts of copper oxalate, 27.7 parts of naphthenoic acid and 120 parts of xylene. The mixture was heated at 120 °C white removing oxalic acid with solvent. A varnish having 51.3 % solids and a viscosity of Y was obtained.

Clear Film Consumption Test

Each of varnishes of Examples 1-12 and Comparative Examples 1-4 was applied on a test panel to a dry film thickness of about 200 μ m. The test panel was attached to a rotating drum tester and rotated continuously at a constant speed (about 15 knot) in the sea water (temperature = 18-23 °C) for 3 months. The film thickness was measured before and after the test. The results are shown in Table 1.

Table 1

Example	Initial film thickness, μm	Film thickness after 3 months, µm	Comsumed film thick- ness, μm
1	193	159	34
2	187	155	32
3	212	174	38
4	197	169	28
5	218	178	40
6	189	156	33
7	175	129	46
8	199	174	25
9	194	163	31
10	223	193	30
11	219	177	42
12	201	166	35
Comp.Ex. 1	189	114	75
Comp.Ex. 2	218	160	58
Comp.Ex. 3	203	140	63
Comp.Ex. 4	184	139	45

Examples 13-24 and Comparative Examples 5-8

Various paint compositions were prepared by milling 15 parts as solids of varnishes of Examples 1-12 and Com-

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parative Examples 1.4. 45 parts of numbers of xide. 4 parts of red nonlock dell 1 parts of tanium disk dell 4 parts of chiprinated paraffin. 1.5 parts of organopentonite and 24.5 parts of xylene. 100 parts in totallifor 5 hours in a parimit Examples 13-24 correspond to Examples 1-12 and Comparative Examples 5-8 correspond to Comparative Examples 1-4 respectively. In Example 20 and Comparative Example 6 copper incodingle was replaced for our dust of xide in Example 15 and Comparative Example 1.3 parts of 2.5 d on pro-2-n-poty. 3- soth azilione were incorporated into the paint formulation as an additional antifolion agent.

Em Consumption Test

Each of paints of Examples 13-24 and Comparative Examples 5-3 was applied on a test panel to a dry film thickness of about 200 km. The test panel was attached to a rotating drum tester and rotated continuously at a constant speed rabout 15 knoth in the sea water, température = 18-23. (C) for 3 months. The film thickness was measured before and after the test. The results are shown in Table 2.

Table 2

Example	nta tim thickness Jim	Film thickness after 3 months	Comsumed film am thickness, am
13	215	15(1	65
- 4	207	147	60
• c ,	196	146	50
* F	•80	135	54
• 7	203	145	55
18	197	148	49
19	199	119	7C
20	205	165	40
ā ·	195	150	45
2 e -	225	172	49
Zn.	,_`1 ¬	1,5,9	7.5
24	198	134	64
Comp Ex 5	193	161	32
Comp Ex 6	219	194	25
Comp Ex 1	334	:00	3.5
្សែកាន្ត 🗜 🖰	•ag	155	3.2

Film integrity Admedich Evaluation

Integ ty evalurion

Earn of paints of Example 19 C4 and Tomparative Examples 5.9 was applied twice using a paint brush to a dry film to exhess of about 100 million onto a steel plate their flues, subjected to a sand blast treatment and then coating of a sustrinof paint. The fest pane was their immersed in the sea water for 6 months, inseed with water allowed to stand for one whole day and eas cated to the integer, in the paint from Their as a first in a condition to the trivial of a security of 1909 and on 1909 as

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Adhesion evaluation:

Using the same test panel as used in the above test, evalution was conducted for adhesion according the method JIS K 5400 • 8 • 5 • 2 • (grid spacing=2mm, number of grid=25) The results of evaluation in terms of the following score are shown in Table 3.

Score

10

20

25

30

35

40

45

55

- 10: Each scratched line was narrow in width and the both sides thereof remained smooth. Peeling was not seen both at intersections of scratched lines and in individual grids.
 - 8: Slight peeling was found at intersections of scratched lines but not in individual grids. The defective area remained less than 5 % of the total square area.
 - 6: Peeling was found both at intersections and either side of the scrathced lines. The defective area extended to 5-15 % of the total square area.
- Wide peeling was found along scratched lines. The defective area extended to 15-35 % of the total square area.
 - 2: Wider peeling than that of score 4 was found. The defective area extended to 35-65 % of the total square area.
 - 0: The defective area extended to greater than 65 % of the total square area.

Table 3

	idble o	
Example	Integrity	Adhesion
13	(©)	10
14	0	8
15	0	8
16	0	10
17	0	8
18	0	8
19	0	8
20	(Ô)	10
21	0	8
22	0	8
23	0	10
24	0	10
Comp.Ex. 5	Х	2
Comp.Ex. 6	Х	2
Comp.Ex. 7	٤.	4
Comp.Ex. 8	Δ	4

Claims

This COC (ALLE NOTES) 4A1 + A

- 50 1. A hydrolyzable, metal-containing resin comprising a copolymer having a plurality of pendant acid groups bound ionically to a metal atom and a monobasic organic acid also bound to the same metal atom as said pendant acid group, said copolymer consisting, in its free acid form, essentially of
 - (a) from 5 to 70 % by weight of a member selected from the group consisting of a (meth)acrylic acid ester having as the ester residue a branched alkyl of four or more carbon atoms having at least one branch on a carbon atom at second to fourth positions from the distal end of the principal chain, a (meth)acrylic acid ester having as the ester residue a cycloalkyl of six or more carbon atoms, a polyalkylene glycol mono(meth)acrylate, a polyalkylene glycol monoalkyl ether (meth)acrylate and an adduct of 2-hydroxyethyl (meth)acrylate with caprolactone;

- ibila proportion of a polymér zable unsaturated práan clacid monomer por esponding to an abdinumber of the resin from 95 to 350 mg kilom glas solid land.
- is the balance of another bolymer zable heutral monomer
- 2. The hydrolyzable limetal-containing resin according to Claim 1, wherein said metal is copper zinc in one licital trianganese or tell-rum.
- 3. A serf-poishing antifouring paint composition comprising the hydic yzable limetal-containing resin of Claim trian antifouring prigment and or an antifouring agent.
- 4. A self-polishing paint composition comprising the hydrolyzable, metal containing resin of Claim 2, an antifouling pigment and or an antifouling agent.
- The antifouring paint composition asserting to Claim 3, where health antifouring pigment is suprous oxite or septer producted.

INTERNATIONAL SEARCH REPORT International application No. PCT/JP95/01625 CLASSIFICATION OF SUBJECT MATTER Int. C16 C08F8/44, C09D5/16, C09D133/02 According to International Patent Classification (IPC) or to both national classification and IPC FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) Int. C16 C08F8/00, 8/44, C09D5/00, 5/16, C09D133/02 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category* Relevant to claim No. Α JP, 5-163311, A (Dainippon Ink & Chemicals, 1 Inc.), June 29, 1993 (29. 06. 93), Claim (Family: none) Further documents are listed in the continuation of Box C. See patent family annex. later document published after the international filing date or provity date and not in conflict with the application but cited to understand the principle or theory underlying the invention Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be "O" document referring to an oral disclosure, use, exhibition or other means considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report October 16, 1995 (16. 10. 95) November 7, 1995 (07. 11. 95) Name and mailing address of the ISA/ Authorized officer

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